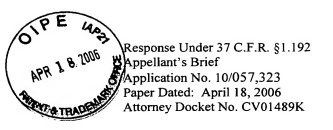
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For FY 2005			Filing Date January 25, 2002					
FOFFY 2005			First Named Inventor Harry R					
Applicant claims	small entity status.	See 37 CFR 1.27	Examiner Name		ng R. Hui			
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Signature	W		(Attorney/Agent) 35,5	712	Telephone 412-47			
Name (Print/Tyne)	Ann Marie Canno	nı			Date April 13	8 2006		

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application of:

Harry R. Davis et al.

: Examiner: San-Ming R. Hui

Serial No.:

10/057,323

Group Art Unit: 1617

Filed: January 25, 2002

: Atty. Docket No.: CV01489K

For:

COMBINATIONS OF PEROXISOME

: Confirmation No. 1525

PROLIFERATOR-ACTIVATED

RECEPTOR (PPAR) ACTIVATOR(S)

AND STEROL ABSORPTION

INHIBITOR(S) AND TREATMENTS

FOR VASCULAR INDICATIONS

MAIL STOP APPEAL BRIEF - PATENTS Commissioner for Patents P.O. Box 1450

Alexandria, VA 22313-1450

ON APPEAL FROM THE PRIMARY EXAMINER TO THE **BOARD OF PATENT APPEALS AND INTERFERENCES**

APPELLANT'S BRIEF UNDER 37 C.F.R. § 1.192

I hereby certify that this correspondence is being deposited with the United States Postal Service with sufficient postage as Express Mail in an envelope addressed to: Commissioner for Patents, P.O. Box 1450 Alexandria, VA 2313-1459

April 18, 2006

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isa Miller

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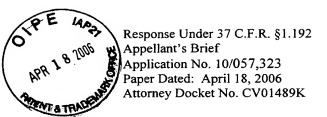
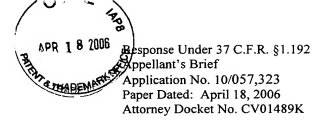


TABLE OF CONTENTS

		Pag	zе
TAB	LE OF	CONTENTS	.I
REA	L PAR	TY IN INTEREST	1
REL	ATED .	APPEALS AND INTERFERENCES	1
STA	TUS O	F CLAIMS	1
STA	rus o	F AMENDMENTS	1
SUM	MARY	OF CLAIMED SUBJECT MATTER	2
GRO	UNDS	OF REJECTION TO BE REVIEWED ON APPEAL:	
I.	1-4, ("Ro	a <u>Prima Facie</u> Case of Obviousness Under 35 U.S.C. § 103 Over Claim 11-13, 37-40, 42, 43, 47-48, 83-84 and 86 as Obvious Over US 5,846,96 senblum et al.") and The Medical Letter on Drugs and Therapeutic 8) 40:1030: 68-69 ("Medical Letter") Been Established?	66 es
ARG	UMEN	T	7
	A.	The Rejection	7
	B.	The Prior Art	7
	C.	Discussion	8
II.	US : Thera Basic	Required Prima Facie Case of Obviousness Under 35 U.S.C. § 103 Ove 5,846,966 ("Rosenblum et al.") and The Medical Letter on Drugs an apeutics (1998) 40:1030: 68-69 ("Medical Letter"), further in view of & Clinical Pharma., 6 th Ed. (1995) 529 ("Katzung") has Failed to bolished	id of oe
	A.	The Rejection10	С
	B.	The Prior Art11	1
	C.	Discussion	1
III.	US 5	Required <u>Prima Facie</u> Case of Obviousness Under 35 U.S.C. § 103 Ove 5,846,966 ("Rosenblum et al.") and Basic & Clinical Pharma., 6 th Ec 5) 529 ("Katzung") has Failed to be Established	d.
	A.	The Rejection13	3
	B.	The Prior Art12	1
	C	Discussion 14	1

I

CLAIMS APPENDIX	16
EVIDENCE APPENDIX	31
RELATED PROCEEDINGS APPENDIX	32



I

REAL PARTY IN INTEREST

The real party in interest in this Appeal is assignee Schering Corporation, having a principal place of business 2000 Galloping Hill Road, Kenilworth, NJ 07033.

II

RELATED APPEALS AND INTERFERENCES

As the legal representative of Appellant, the undersigned attorney has no knowledge of any appeals or interferences directly related to this Appeal.

Ш

STATUS OF CLAIMS

This is an original patent application in which claims 1-4, 11-13, 21, 28, 32, 34, 37-40, 42, 43, 47, 48, 83, 84, 86, 100 and 101 are pending in the application. Claims 5-10, 14-20, 22-31, 33, 35, 36, 41, 44-46, 49-82, 85 and 87-99 have been withdrawn from consideration by the Examiner as being non-elected.

Claims 1-4, 11-13, 21, 28, 32, 34, 37-40, 42, 43, 47, 48, 83, 84, 86, 100 and 101 (pending) were finally rejected under 35 U.S.C. §103(a) in an Office Action mailed November 22, 2005 ("Final Office Action") and Advisory Action mailed April 10, 2006 ("Advisory Action").

Twenty-four (24) pending claims (1-4, 11-13, 21, 28, 32, 34, 37-40, 42, 43, 47, 48, 83, 84, 86, 100 and 101) are at issue in this Appeal.

IV

STATUS OF AMENDMENTS

No claims were amended after final rejection. A copy of the claims involved in this Appeal is contained in the Appendix attached hereto.

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

V

SUMMARY OF CLAIMED SUBJECT MATTER

In embodiments set forth in claim 1, Applicants have discovered a composition comprising:

- (a) at least one peroxisome proliferator-activated receptor (PPAR) activator; and
- (b) at least one sterol absorption inhibitor represented by Formula (I):

$$Ar^{1}-X_{m}-(C)_{q}-Y_{n}-(C)_{r}-Z_{p}$$
 Ar^{3}
 Ar^{2}
 Ar^{2}

(I)

or isomers thereof, or pharmaceutically acceptable salts, solvates or prodrugs thereof (see original claim 1 for moiety definitions). See original claim 1 and page 3, line 6 - page 4, line 17 of the specification.

In another embodiment set forth in Claim 37, Applicants have discovered a therapeutic combination comprising:

- (a) a first amount of at least one peroxisome proliferator-activated receptor activator; and
- (b) a second amount of at least one sterol absorption inhibitor represented by Formula (I):

$$Ar^{1}-X_{m}-(C)_{q}-Y_{n}-(C)_{r}-Z_{p}$$
 Ar^{3}
 Ar^{2}

(I)

or isomers thereof, or pharmaceutically acceptable salts, solvates or prodrugs thereof (see original claim 37 for moiety definitions). See original claim 37 and page 21, line 27 - page 22, line 7 of the specification.

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

In another embodiment set forth in Claim 42, Applicants have discovered a composition comprising:

- (a) at least one fibric acid derivative; and
- (b) a compound represented by Formula (II) below:

or pharmaceutically acceptable salt or solvate thereof, or prodrug of the compound of Formula (II) or of the salt or solvate thereof. <u>See</u> original claim 42 and page 4, lines 18-22 of the specification.

In another embodiment set forth in Claim 48, Applicants have discovered a therapeutic combination comprising:

- (a) a first amount of at least one fibric acid derivative; and
- (b) a second amount of a compound represented by Formula (II) below:

or pharmaceutically acceptable salt or solvate thereof, or prodrug of the compound of Formula (II) or of the salt or solvate thereof, wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment or prevention of a vascular condition, diabetes, obesity or lowering a concentration of a

sterol in plasma of a mammal. See original claim 48 and page 4, lines 18-22 of the specification.

In another embodiment set forth in Claim 83, Applicants have discovered a composition comprising:

- (a) at least one peroxisome proliferator-activated receptor activator; and
- (b) at least one sterol absorption inhibitor represented by Formula (IX):

$$Ar^1$$
 CH Q R_{26} R_{26} R_{26}

or isomers thereof, or pharmaceutically acceptable salts, solvates or prodrugs thereof (see original claim 83 for moiety definitions). See original claim 83 and page 18, line 24 - page 21, line 26 of the specification.

In another embodiment set forth in Claim 86, Applicants have discovered a therapeutic combination comprising:

- (a) a first amount of at least one peroxisome proliferator-activated receptor activator; and
- (b) a second amount of at least one sterol absorption inhibitor represented by Formula (IX):

$$Ar^1$$
— CH — Q — R_{26}
 Ar^2
(IX)

or isomers thereof, or pharmaceutically acceptable salts, solvates or prodrugs thereof, wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment or prevention of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal. (see original claim 86 for moiety definitions). See original claim 86 and page 4, lines 18-22 of the specification.

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

In another embodiment set forth in Claim 100, Applicants have discovered a composition comprising (a) at least one antioxidant or vitamin and (b) at least one substituted azetidinone compound or substituted β-lactam compound or isomers thereof, pharmaceutically acceptable salts or solvates, or prodrugs thereof. See original claim 100 at page 162, lines 1-7 of the specification.

In another embodiment set forth in Claim 101, Applicants have discovered a therapeutic combination comprising (a) a first amount of at least one antioxidant or vitamin and (b) a second amount of at least one substituted azetidinone compound or substituted β-lactam compound or isomers thereof, or pharmaceutically acceptable salts or solvates of the at least one substituted azetidinone compound or the at least one substituted β-lactam compound compound or isomers thereof, pharmaceutically acceptable salts or solvates, or prodrugs thereof, wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal. See original claim 101 at page 162, lines 7-18 of the specification.

In the Office Action of July 2, 2003, Applicants were required to elect a species of peroxisome proliferator-activated receptor (PPAR) activator, sterol absorption inhibitor, and third therapeutic agent.

Applicants provisionally elected with traverse fenofibrate as the PPAR activator. See Response to Restriction Requirement and Election of Species of August 1, 2003 ("Response") at page 2, lines 8-9.

Applicants provisionally elected with traverse ezetimibe as the sterol absorption inhibitor, represented by Formula (II) below:

Ezetimibe is the active ingredient in ZETIATM (ezetimibe) pharmaceutical formulation and VYTORINTM (ezetimibe/simvastatin) pharmaceutical formulation, both of which are commercially available from MSP (Merck Schering-Plough) Pharmaceuticals, Inc. See Response to Restriction Requirement and Election of Species of August 1, 2003 ("Response") at page 2, lines 12-13.

In the same Response, Applicants provisionally elected niacin as the third therapeutic agent. See Response to Restriction Requirement and Election of Species of August 1, 2003 ("Response") at page 2, lines 15-16.

The claimed compositions and combinations can be useful for treating vascular conditions, diabetes, obesity and/or lowering concentration of a sterol in plasma in a mammal (page 22, lines 8-15 of the specification).

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

\mathbf{VI}

GROUNDS OF REJECTION TO BE REVIEWED ON APPEAL

I. Has a <u>Prima Facie</u> Case of Obviousness Under 35 U.S.C. § 103 Over Claims 1-4, 11-13, 37-40, 42, 43, 47-48, 83-84 and 86 as Obvious Over US 5,846,966 ("Rosenblum et al.") and The Medical Letter on Drugs and Therapeutics (1998) 40:1030: 68-69 ("Medical Letter") Been Established?

VII

<u>ARGUMENT</u>

A. The Rejection

Claims 1-4, 11-13, 37-40, 42, 43, 47-48, 83-84 and 86 have been rejected under 35 U.S.C. §103(a) as obvious over US 5,846,966 ("Rosenblum et al.") and The Medical Letter on Drugs and Therapeutics (1998) 40:1030: 68-69 ("Medical Letter").

The reasons for rejection are set forth in the Final Office Action of November 22, 2005, summarized as follows:

Rosenblum et al. disclose that the elected compound of Formula II, ezetimibe, is useful for reducing cholesterol and the risk of atherosclerosis (Office Action at page 4). Medical Letter teaches fenofibrate as useful in reducing serum cholesterol (Final Office Action at page 4).

It is acknowledged in the Final Office Action that the primary references do not expressly teach the claimed composition comprising ezetimibe and fenofibrate (Final Office Action at page 4).

It is alleged that it would have been obvious to one of ordinary skill in the art at the time the invention was made to combine ezetimibe and fenofibrate, since the cited prior art teaches that both ezetimibe and fenofibrate are useful in reducing serum cholesterol individually, citing *In re Kerkoven*, 205 U.S.P.Q. 1069 (Final Office Action at pages 4-5).

B. The Prior Art

Rosenblum et al. disclose the compound of Formula II (Page 29, Ex. 6). Rosenblum et al. disclose starch-based pharmaceutical compositions including compounds of Formula I of Rosenblum et al. (Ex. A and B, Page 29). Rosenblum et al. teach that the active compounds therein can be combined with HMG CoA

reductase inhibitors, such as simvastatin (Page 5, paragraph 0028). Rosenblum et al. also disclose that the active compounds are useful for reducing cholesterol and the risk of atherosclerosis (claims).

Medical Letter teaches fenofibrate as useful in reducing VLDL cholesterol and triglycerides (Medical Letter at page 68).

C. The Required Prima Facie Case of Obviousness Under 35 U.S.C. § 103 Has Not Been Established

When making a rejection under 35 U.S.C. § 103, the Examiner has the burden of establishing a <u>prima facie</u> case of obviousness. <u>In re Fritch</u>, 23 U.S.P.Q.2d 1780, 1783 (Fed. Cir. 1992).

The Examiner can satisfy this burden only by showing an objective teaching in the prior art, or knowledge generally available to one of ordinary skill in the art, which would lead an individual to combine the relevant teachings of the references [and/or the knowledge] in the manner suggested by the Examiner. <u>Id.</u>; <u>In re Fine</u>, 5 U.S.P.Q.2d 1596, 1598 (Fed. Cir. 1988).

The mere fact that the prior art could be modified does not make the modification obvious unless the prior art suggests the desirability of the modification (emphasis added). <u>In re Fritch</u>, 23 U.S.P.Q.2d at 1784; <u>In re Laskowski</u>, 10 U.S.P.Q.2d 1397, 1398 (Fed. Cir. 1989); <u>In re Gordon</u>, 221 U.S.P.Q. 1125, 1127 (Fed. Cir. 1984).

"The ultimate determination of patentability must be based on consideration of the entire record, by a preponderance of evidence, with due consideration to the persuasiveness of any arguments and any secondary evidence." Manual of Patent Examining Procedure, (Rev. 1, Feb. 2003) § 716.01(d) and In re Oetiker, 24 U.S.P.Q.2d 1443, 1444 (Fed. Cir. 1992).

Claims 1 and 37 recite a composition and therapeutic combination, respectively, comprising a sterol absorption inhibitor of Formula I shown above, isomers, prodrugs, or pharmaceutically acceptable salts or solvates thereof; and at least one PPAR activator.

Claims 2 and 38 depend from claims 1 and 37, respectively, and recite that the at least one PPAR activator is a fibric acid derivative.

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006

Attorney Docket No. CV01489K

Claim 3 depends from claim 2 and recites that the fibric acid derivative is selected from, *inter alia*, fenofibrate. Claim 4 depends from claim 3 and recites that the fibric acid derivative is fenofibrate.

Claim 13 depends from claim 1 and recites that the amount of sterol absorption inhibitor administered to a mammal ranges from about 0.1 to about 1000 mg/day.

Claim 39 depends from claim 37 and recites that the PPAR activator is administered concomitantly with the sterol absorption inhibitor.

Claim 40 depends from claim 37 and recites that the PPAR activator and the sterol absorption inhibitor are present in separate treatment compositions.

Claims 42 and 48 recite a composition and therapeutic combination, respectively, comprising ezetimibe and at least one fibric acid derivative.

Claim 43 depends from claim 42 and recites that the fibric acid derivative is fenofibrate.

Claim 47 recites a pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal using the composition of claim 42 and carrier.

Claims 83 and 86 recite a composition and therapeutic combination, respectively, comprising a sterol absorption inhibitor of Formula IX shown above, isomers, prodrugs, or pharmaceutically acceptable salts or solvates thereof; and at least one PPAR activator.

Claim 84 pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal using the composition of claim 83 and carrier.

It is respectfully submitted that the combination of the references cited as rendering the claimed invention obvious is improper because there is no suggestion in the cited references to combine the claimed components of sterol absorption inhibitor (such as that of Formula (I) (e.g., ezetimibe)) and PPAR activator (such as fenofibrate).

Neither Rosenblum et al. nor Medical Letter provides motivation for substituting a PPAR activator for the statin used in combination with ezetimibe described in Rosenblum et al. As disclosed in the Medical Letter Clinical Study

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006

Attorney Docket No. CV01489K

section at page 68, fenofibrate is not as effective as statins in lowering LDL cholesterol, a major risk factor in atherogenesis. Since statins are more effective in lowering LDL cholesterol, there is no motivation to substitute a PPAR activator such as fenofibrate for the statin in the combination disclosed in Rosenblum et al.

There is no guidance provided by Rosenblum et al. nor Medical Letter to pick and choose among numerous cholesterol treatments to select the particularly claimed combination of sterol absorption inhibitor (such as that of Formula (II) (e.g., ezetimibe)) and PPAR activator (such as fenofibrate).

Therefore, the prima facie case of obviousness based upon Rosenblum et al. and Medical Letter has not been established and the rejection of claims 1-4, 11-13, 37-40, 42, 43, 47, 48, 83, 84 and 86 should be reconsidered and withdrawn.

II. The Required <u>Prima Facie</u> Case of Obviousness Under 35 U.S.C. § 103 Over US 5,846,966 ("Rosenblum et al.") and The Medical Letter on Drugs and Therapeutics (1998) 40:1030: 68-69 ("Medical Letter"), further in view of Basic & Clinical Pharma., 6th Ed. (1995) 529 ("Katzung") has Failed to be Established

A. The Rejection

Claims 21, 28, 32 and 34 were rejected under 35 U.S.C. §103(a) as obvious over Rosenblum et al. and the Medical Letter, further in view of Basic & Clinical Pharma., 6th Ed. (1995) 529 ("Katzung").

The reasons for rejection are set forth in the Final Office Action, summarized as follows:

Rosenblum et al. and Medical Letter suggest a composition containing fenofibrate and ezetimibe (Final Office Action at page 5).

It is acknowledged that the primary references do not expressly teach the claimed composition containing niacin (Final Office Action at page 5).

Katzung teaches niacin as useful for lowering cholesterol (Final Office Action at page 6).

It is alleged that it would have been obvious to one of ordinary skill in the art at the time the invention was made to incorporate niacin into the ezetimibe and fenofibrate composition, since the cited prior art teaches that all three ingredients are

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006

Attorney Docket No. CV01489K

useful in reducing serum cholesterol, citing In re Kerkoven, 205 U.S.P.Q. 1069

(Office Action at page 6).

B. The Prior Art

Rosenblum et al. disclose the compound of Formula II (Page 29, Ex. 6).

Rosenblum et al. disclose starch-based pharmaceutical compositions including

compounds of Formula I of Rosenblum et al. (Ex. A and B, Page 29). Rosenblum et

al. teach that the active compounds therein can be combined with HMG CoA

reductase inhibitors, such as simvastatin (Page 5, paragraph 0028). Rosenblum et al.

also disclose that the active compounds are useful for reducing cholesterol and the

risk of atherosclerosis (claims). Rosenblum et al. do not disclose niacin.

Medical Letter teaches fenofibrate as useful in reducing VLDL cholesterol and

triglycerides (Medical Letter at page 68). Medical Letter discloses that niacin is a

drug for treating hypertriglyceridemia (Medical Letter at page 69). Medical Letter

does not suggest or disclose a combination of substituted azetidinone compound,

PPAR activator and niacin.

Katzung discloses that niacin decreases VLDL and LDL levels in patients

(Katzung at 529). Katzung does not suggest or disclose a combination of substituted

azetidinone compound, PPAR activator and niacin.

C. The Required Prima Facie Case of Obviousness Under 35 U.S.C. § 103

Has Not Been Established

Claim 21 depends from claim 1 and recite that the composition further

comprises nicotinic acid, niceritrol, nicofuranose or acipimox. Claim 28 depends

from claim 1 and recites that the composition further comprises at least one

antioxidant or vitamin. Thus the composition would comprise sterol absorption

inhibitor, PPAR activator such as fenofibrate, and niacin, for example.

Claim 32 depends from claim 1 and recites that the composition further

comprises at least one cardiovascular agent selected from the group consisting of

calcium channel blockers, adrenergic blockers, adrenergic stimulants, angiotensin

converting enzyme (ACE) inhibitors, antihypertensive, angiotensin II receptor

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006

Attorney Docket No. CV01489K

antagonists, anti-anginal agents, coronary vasodilators, diuretics and combinations thereof.

Claim 34 recites a pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal using the composition of claim 1 and carrier. Claim 34 does not require the presence of nicotinic acid, niceritrol, nicofuranose or acipimox, although such compounds could be present.

With respect to claims 21 and 28, Rosenblum et al. nor Medical Letter, taken alone or together as suggested in the Office Action, provides any motivation for a triple combination treatment of sterol absorption inhibitor (such as that of Formula (II) (e.g., ezetimibe)), PPAR activator (such as fenofibrate) and niacin. references provide no guidance or motivation as to the desirability for such as combination or selecting the particular components of the combination, or the potential effect of drug-drug interactions. For example, in the Drug Interaction section at page 69, Medical Letter discloses that it is unclear whether, like gemfibrozil and niacin, concurrent administration of fenofibrate with a statin could increase the risk of rhabdomyolysis. In the Advisory Action of December 7, 2004, the Examiner encouraged Applicants to bring forth evidence of potential drug-drug interaction. This evidence is present in the Drug Interaction section at page 69 of Medical Letter cited in the rejection as pointed out above and the burden therefore is shifted to the Examiner to refute the teaching in the reference which was cited in the rejection. Applicants have not mischaracterized the teachings of this reference as alleged in the Final Office Action at page 6. One skilled in the art would consider such a statement regarding the potential for drug interaction worthy of further serious consideration.

Katzung provides no further incentive to one skilled in the art to include niacin in a composition or therapeutic combination of sterol absorption inhibitor and PPAR activator.

Because of the difference of the way that each component of the presently claimed combination acts, it is respectfully submitted that the rejection is based upon an improper combination of references.

With respect to claim 32, Rosenblum et al., Medical Letter, nor Katzung, taken alone or together as suggested in the Office Action, provides any motivation for

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12

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006

Attorney Docket No. CV01489K

a triple combination treatment of sterol absorption inhibitor (such as that of Formula (II) (e.g., ezetimibe)), PPAR activator (such as fenofibrate) and at least one cardiovascular agent selected from the group consisting of calcium channel blockers, adrenergic blockers, adrenergic stimulants, angiotensin converting enzyme (ACE) inhibitors, antihypertensive, angiotensin II receptor antagonists, anti-anginal agents, coronary vasodilators, diuretics and combinations thereof.

With respect to claim 34, Rosenblum et al., Medical Letter, nor Katzung, taken alone or together as suggested in the Office Action, provides any motivation for a pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal using the composition of claim 1 and carrier. Claim 34 does not require the presence of nicotinic acid, niceritrol, nicofuranose or acipimox, although such compounds could be present.

Therefore, the prima facie case of obviousness based upon Rosenblum et al., Medical Letter and Katzung has not been established and the rejection of claims 21, 28, 32 and 34 should be reconsidered and withdrawn.

III. The Required Prima Facie Case of Obviousness Under 35 U.S.C. § 103 Over US 5,846,966 ("Rosenblum et al.") and Basic & Clinical Pharma., 6th Ed. (1995) 529 ("Katzung") has Failed to be Established

A. The Rejection

Claims 100 and 101 were rejected under 35 U.S.C. §103(a) as obvious over Rosenblum et al. and Basic & Clinical Pharma., 6th Ed. (1995) 529 ("Katzung").

The reasons for rejection are set forth in the Final Office Action, summarized as follows:

Rosenblum et al. teaches that ezetimibe is useful for reducing cholesterol and the risk of atherosclerosis (Final Office Action at page 7).

Katzung teaches niacin as useful for lowering cholesterol (Final Office Action at pages 7-8).

It is acknowledged that the primary references do not expressly teach the claimed composition containing niacin (Final Office Action at page 8).

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006

Attorney Docket No. CV01489K

It is alleged that it would have been obvious to one of ordinary skill in the art

at the time the invention was made to incorporate niacin into the ezetimibe

composition, since the cited prior art teaches that both ingredients are useful in

reducing serum cholesterol, citing In re Kerkoven, 205 U.S.P.Q. 1069 (Final Office

Action at page 8).

B. The Prior Art

Rosenblum et al. disclose the compound of Formula II (Page 29, Ex. 6).

Rosenblum et al. disclose starch-based pharmaceutical compositions including

compounds of Formula I of Rosenblum et al. (Ex. A and B Page 29). Rosenblum et

al. teach that the active compounds therein can be combined with HMG CoA

reductase inhibitors, such as simvastatin (Page 5, paragraph 0028). Rosenblum et al.

also disclose that the active compounds are useful for reducing cholesterol and the

risk of atherosclerosis (claims). Rosenblum et al. do not disclose niacin.

Katzung discloses that niacin decreases VLDL and LDL levels in patients

(Katzung at 529). Katzung does not suggest or disclose a combination of substituted

azetidinone compound and niacin.

C. The Required Prima Facie Case of Obviousness Under 35 U.S.C. § 103

Has Not Been Established

Claims 100 and 101 recite a composition or therapeutic combination

comprising (a) at least one antioxidant or vitamin and (b) at least one substituted

azetidinone compound or substituted β-lactam compound or isomers, prodrugs, salts

or solvates thereof.

With respect to patentability of the composition or combination of Claims 100

and 101, neither Rosenblum nor Katzung suggests or disclose combinations of a sterol

absorption inhibitor and antioxidant or vitamin.

Therefore, the prima facie case of obviousness based upon Rosenblum et al.

and Katzung has not been established and the rejection of claims 100 and 101 should

be reconsidered and withdrawn.

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14

Accordingly, Applicants respectfully request that the § 103(a) rejections of claims 1-4, 11-13, 21, 28, 32, 34, 37-40, 42, 43, 47-48, 83, 84, 86 and 100-101 be reconsidered and withdrawn.

Respectfully submitted,

Date: April 18, 2006

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Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

CLAIM APPENDIX

- 1. A composition comprising:
- (a) at least one peroxisome proliferator-activated receptor activator; and
- (b) at least one sterol absorption inhibitor represented by Formula (I):

$$Ar^{1}-X_{m}-(C)_{q}-Y_{n}-(C)_{r}-Z_{p}$$
 Ar^{3}
 Ar^{2}
 Ar^{2}

(I)

or isomers thereof, or pharmaceutically acceptable salts or solvates of the compounds of Formula (I) or of the isomers thereof, or prodrugs of the compounds of Formula (I) or of the isomers, salts or solvates thereof,

wherein in Formula (I) above:

 Ar^{1} and Ar^{2} are independently selected from the group consisting of aryl and R^{4} -substituted aryl;

Ar³ is aryl or R⁵-substituted aryl;

X, Y and Z are independently selected from the group consisting of -CH₂-, -CH(lower alkyl)- and -C(dilower alkyl)-;

R and R² are independently selected from the group consisting of -OR⁶, -O(CO)R⁶, -O(CO)OR⁹ and -O(CO)NR⁶R⁷;

 R^{1} and R^{3} are independently selected from the group consisting of hydrogen, lower alkyl and aryl;

q is 0 or 1;

r is 0 or 1;

m, n and p are independently selected from 0, 1, 2, 3 or 4; provided that at least one of q and r is 1, and the sum of m, n, p, q and r is 1, 2, 3, 4, 5 or 6; and provided that when p is 0 and r is 1, the sum of m, q and n is 1, 2, 3, 4 or 5;

 R^4 is 1-5 substituents independently selected from the group consisting of lower alkyl, $-OR^6$, $-O(CO)R^6$, $-O(CO)OR^9$, $-O(CH_2)_{1-5}OR^6$, $-O(CO)NR^6R^7$, $-NR^6R^7$, $-NR^6(CO)R^7$, $-NR^6(CO)OR^9$, $-NR^6(CO)NR^7R^8$, $-NR^6SO_2R^9$, $-COOR^6$, $-CONR^6R^7$, $-COR^6$, $-SO_2NR^6R^7$, $S(O)_{0-2}R^9$, $-O(CH_2)_{1-10}$ - $-COOR^6$, $-O(CH_2)_{1-10}CONR^6R^7$, $-(lower alkylene)COOR^6$, $-CH=CH-COOR^6$, $-CF_3$, -CN, $-NO_2$ and halogen;

 R^5 is 1-5 substituents independently selected from the group consisting of $-OR^6$, $-O(CO)R^6$, $-O(CO)OR^9$, $-O(CH_2)_{1-5}OR^6$, $-O(CO)NR^6R^7$, $-NR^6R^7$, $-NR^6(CO)R^7$, $-NR^6(CO)OR^9$, $-NR^6(CO)NR^7R^8$, $-NR^6SO_2R^9$, $-COOR^6$, $-CONR^6R^7$, $-COR^6$, $-SO_2NR^6R^7$, $S(O)_{0-2}R^9$, $-O(CH_2)_{1-10}$ - $-COOR^6$, $-O(CH_2)_{1-10}CONR^6R^7$, $-(lower alkylene)COOR^6$ and $-CH=CH-COOR^6$;

R⁶, R⁷ and R⁸ are independently selected from the group consisting of hydrogen, lower alkyl, aryl and aryl-substituted lower alkyl; and R⁹ is lower alkyl, aryl or aryl-substituted lower alkyl.

- 2. The composition according to claim 1, wherein the at least one peroxisome proliferator-activated receptor activator is a fibric acid derivative.
- 3. The composition according to claim 2, wherein the fibric acid derivative is selected from the group consisting of fenofibrate, clofibrate, gemfibrozil, ciprofibrate, bezafibrate, clinofibrate, binifibrate, lifibrol and mixtures thereof.
- 4. The composition according to claim 3, wherein the fibric acid derivative comprises fenofibrate.
- 11. The composition according to claim 1, wherein the at least one peroxisome proliferator-activated receptor activator is administered to a mammal in an amount ranging from about 50 to about 3000 milligrams of peroxisome proliferator-activated receptor activator per day.

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

12. The composition according to claim 1, wherein the sterol absorption inhibitor is represented by Formula (II) below:

or pharmaceutically acceptable salt or solvate thereof, or prodrug of the compound of Formula (II) or of the salt or solvate thereof.

- 13. The composition according to claim 1, wherein the at least one sterol absorption inhibitor is administered to a mammal in an amount ranging from about 0.1 to about 1000 milligrams of sterol absorption inhibitor per day.
- 21. The composition according to claim 1, further comprising nicotinic acid, niceritrol, nicofuranose or acipimox.
- 28. The composition according to claim 1, further comprising at least one antioxidant or vitamin.
- 32. The composition according to claim 1, further comprising at least one cardiovascular agent selected from the group consisting of calcium channel blockers, adrenergic blockers, adrenergic stimulants, angiotensin converting enzyme (ACE)

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

inhibitors, antihypertensive, angiotensin II receptor antagonists, anti-anginal agents, coronary vasodilators, diuretics and combinations thereof.

- 34. A pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal, comprising a therapeutically effective amount of the composition of claim 1 and a pharmaceutically acceptable carrier.
 - 37. A therapeutic combination comprising:
 - (a) a first amount of at least one peroxisome proliferator-activated receptor activator; and
 - (b) a second amount of at least one sterol absorption inhibitor represented by Formula (I):

$$R = R^{2}$$
 $Ar^{1}-X_{m}-(C)_{q}-Y_{n}-(C)_{r}-Z_{p}$
 $R^{1} = R^{3}$
 Ar^{2}

(I)

or isomers thereof, or pharmaceutically acceptable salts or solvates of the compounds of Formula (I) or of the isomers thereof, or prodrugs of the compounds of Formula (I) or of the isomers, salts or solvates thereof, wherein:

 Ar^{1} and Ar^{2} are independently selected from the group consisting of aryl and R^{4} -substituted aryl;

Ar is aryl or R -substituted aryl;

X, Y and Z are independently selected from the group consisting of

-CH2-, -CH(lower alkyl)- and -C(dilower alkyl)-;

R and R² are independently selected from the group consisting of -OR⁶, -O(CO)R⁶, -O(CO)OR⁹ and -O(CO)NR⁶R⁷;

R¹ and R³ are independently selected from the group consisting of hydrogen, lower alkyl and aryl;

q is 0 or 1;

r is 0 or 1;

m, n and p are independently selected from 0, 1, 2, 3 or 4; provided that at least one of q and r is 1, and the sum of m, n, p, q and r is 1, 2, 3, 4, 5 or 6; and provided that when p is 0 and r is 1, the sum of m, q and n is 1, 2, 3, 4 or 5;

 R^4 is 1-5 substituents independently selected from the group consisting of lower alkyl, $-OR^6$, $-O(CO)R^6$, $-O(CO)OR^9$, $-O(CH_2)_{1-5}OR^6$, $-O(CO)NR^6R^7$, $-NR^6R^7$, $-NR^6(CO)R^7$, $-NR^6(CO)OR^9$, $-NR^6(CO)NR^7R^8$, $-NR^6SO_2R^9$, $-COOR^6$, $-CONR^6R^7$, $-COR^6$, $-SO_2NR^6R^7$, $S(O)_{0-2}R^9$, $-O(CH_2)_{1-10}$ - $-COOR^6$, $-O(CH_2)_{1-10}CONR^6R^7$, $-(lower alkylene)COOR^6$, $-CH=CH-COOR^6$, $-CF_3$, -CN, $-NO_2$ and halogen;

 R^5 is 1-5 substituents independently selected from the group consisting of $-OR^6$, $-O(CO)R^6$, $-O(CO)OR^9$, $-O(CH_2)_{1-5}OR^6$, $-O(CO)NR^6R^7$, $-NR^6R^7$, $-NR^6(CO)R^7$, $-NR^6(CO)OR^9$, $-NR^6(CO)NR^7R^8$, $-NR^6SO_2R^9$, $-COOR^6$, $-CONR^6R^7$, $-COR^6$, $-SO_2NR^6R^7$, $S(O)_{0-2}R^9$, $-O(CH_2)_{1-10}$ - $-COOR^6$, $-O(CH_2)_{1-10}CONR^6R^7$, $-(lower alkylene)COOR^6$ and $-CH=CH-COOR^6$;

R⁶, R⁷ and R⁸ are independently selected from the group consisting of hydrogen, lower alkyl, aryl and aryl-substituted lower alkyl; and

R⁹ is lower alkyl, aryl or aryl-substituted lower alkyl wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal.

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

- 38. A therapeutic combination according to claim 37, wherein the at least one peroxisome proliferator-activated receptor activator is a fibric acid derivative.
- 39. A therapeutic combination according to claim 37, wherein the at least one peroxisome proliferator-activated receptor activator is administered concomitantly with the at least one sterol absorption inhibitor.
- 40. A therapeutic combination according to claim 37, wherein the at least one peroxisome proliferator-activated receptor activator and the at least one sterol absorption inhibitor are present in separate treatment compositions.
 - 42. A composition comprising:
 - (a) at least one fibric acid derivative; and
 - (b) a compound represented by Formula (II) below:

or pharmaceutically acceptable salt or solvate thereof, or prodrug of the compound of Formula (II) or of the salt or solvate thereof.

43. The composition according to claim 42, wherein the fibric acid derivative is fenofibrate.

Response Under 37 C.F.R. §1.192 Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

- 47. A pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal, comprising a therapeutically effective amount of the composition of claim 42 and a pharmaceutically acceptable carrier.
 - 48. A therapeutic combination comprising:
 - (a) a first amount of at least one fibric acid derivative; and
 - (b) a second amount of a compound represented by Formula (II) below:

or pharmaceutically acceptable salt or solvate thereof, or prodrug of the compound of Formula (II) or of the salt or solvate thereof,

wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal.

- 83. A composition comprising:
- (a) at least one peroxisome proliferator-activated receptor activator; and
- (b) at least one sterol absorption inhibitor represented by Formula (IX):

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

$$Ar^1$$
— CH — Q — R_{26}
 Ar^2
(IX)

or isomers thereof, or pharmaceutically acceptable salts or solvates of the compounds of Formula (IX) or of the isomers thereof, or prodrugs of the compounds of Formula (IX) or of the isomers, salts or solvates thereof, wherein, in Formula (IX) above,

R²⁶ is selected from the group consisting of:

- a) OH;
- b) OCH₃;
- c) fluorine and
- d) chlorine;

R¹ is selected from the group consisting of

H,
$$OR^5$$
 OR^4 OR^7 OR^7 OR^7 OR^7 OR^7 OR^7 OR^8 OR^8

R, R^a and R^b are independently selected from the group consisting of H, -OH, halogeno, -NH₂, azido, (C₁-C₆)alkoxy(C₁-C₆)-alkoxy and -W-R³⁰;

W is independently selected from the group consisting of -NH-C(O)-, -O-C(O)-, $-O-C(O)-N(R^{31})-$, $-NH-C(O)-N(R^{31})-$ and $-O-C(S)-N(R^{31})-$;

 R^2 and R^6 are independently selected from the group consisting of H, (C1-C6)alkyl, aryl and aryl(C1-C6)alkyl;

 R^3 , R^4 , R^5 , R^7 , R^{3a} and R^{4a} are independently selected from the group consisting of H, (C1-C6)alkyl, aryl(C1-C6)alkyl, -C(O)(C1-C6)alkyl and -C(O)aryl;

 R^{30} is independently selected form the group consisting of R^{32} -substituted T, R^{32} -substituted-T-(C₁-C₆)alkyl, R^{32} -substituted-(C₂-C₄)alkenyl, R^{32} -substituted-(C₃-C₇)cycloalkyl, R^{32} -substituted-(C₃-C₇)cycloalkyl(C₁-C₆)alkyl;

R³¹ is independently selected from the group consisting of H and (C₁-C₄)alkyl;

T is independently selected from the group consisting of phenyl, furyl, thienyl, pyrrolyl, oxazolyl, isoxazolyl, thiazolyl, iosthiazolyl, benzothiazolyl, thiadiazolyl, pyrazolyl, imidazolyl and pyridyl;

R³² is independently selected from 1-3 substituents independently selected from the group consisting of H, halogeno, (C₁-C₄)alkyl, -OH, phenoxy, -CF₃, -NO₂, (C₁-C₄)alkoxy, methylenedioxy, oxo, (C₁-C₄)alkylsulfanyl, (C₁-C₄)alkylsulfinyl, (C₁-C₄)alkylsulfonyl, -N(CH₃)₂, -C(O)-NH(C₁-C₄)alkyl, -C(O)-N((C₁-C₄)alkyl)₂, -C(O)-(C₁-C₄)alkyl, -C(O)-(C₁-C₄)alkoxy and pyrrolidinylcarbonyl; or R³² is a covalent bond and R³¹, the nitrogen to which it is attached and R³² form a pyrrolidinyl, piperidinyl, N-methyl-piperazinyl, indolinyl or morpholinyl group, or a (C₁-C₄)alkoxycarbonyl-substituted pyrrolidinyl, piperidinyl, N-methylpiperazinyl, indolinyl or morpholinyl group;

Ar¹ is aryl, R¹⁰-substituted aryl; pyridyl, isoxazolyl, furanyl, pyrrolyl, thienyl, imidazolyl, pyrazolyl, thiazolyl, pyrazinyl, pyrimidinyl or pyridazinyl;

Ar² is aryl or R¹¹-substituted aryl;

Q is $-(CH_2)_q$ -, wherein q is 2-6, or, with the 3-position ring carbon of the azetidinone,

$$R^{12} - (R^{13})_a$$
 forms the spiro group $(R^{14})_b$;

Response Under 37 C.F.R. §1.192 Appellant's Brief Application No. 10/057,323

Application No. 10/05 /,323
Paper Dated: April 18, 2006
Attorney Docket No. CV01489K

 R^{12} is

 R^{13} and R^{14} are independently selected from the group consisting of -CH₂-, -CH(C₁-C₆ alkyl)-, -C(di-(C₁-C₆) alkyl), -CH=CH- and -C(C₁-C₆ alkyl)=CH-; or R^{12} together with an adjacent R^{13} , or R^{12} together with an adjacent R^{14} , form a -CH=CH- or a -CH=C(C₁-C₆ alkyl)- group;

a and b are independently 0, 1, 2 or 3, provided both are not zero; provided that when R^{13} is -CH=CH- or -C(C₁-C₆ alkyl)=CH-, a is 1; provided that when R^{14} is -CH=CH- or -C(C₁-C₆ alkyl)=CH-, b is 1; provided that when a is 2 or 3, the R^{13} 's can be the same or different; and provided that when b is 2 or 3, the R^{14} 's can be the same or different;

 R^{10} and R^{11} are independently selected from the group consisting of 1-3 substituents independently selected from the group consisting of (C₁-C₆)alkyl, -OR¹⁹, -O(CO)R¹⁹, -O(CO)OR²¹, -O(CH₂)₁₋₅OR¹⁹, -O(CO)NR¹⁹R²⁰, -

 $-OR^{19}$, $-O(CO)R^{19}$, $-O(CO)OR^{21}$, $-O(CH_2)_{1-5}OR^{19}$, $-O(CO)NR^{19}R^{20}$, $NR^{19}R^{20}$,

 $-NR^{19}(CO)R^{20}$, $-NR^{19}(CO)OR^{21}$, $-NR^{19}(CO)NR^{20}R^{25}$, $-NR^{19}SO_2R^{21}$, $-COOR^{19}$,

 $-\text{CONR}^{19}\text{R}^{20}, -\text{COR}^{19}, -\text{SO}_2\text{NR}^{19}\text{R}^{20}, -\text{S(O)}_{0\text{-}2}\text{R}^{21}, -\text{O(CH}_2)_{1\text{-}10}\text{-COOR}^{19},$

 $-O(CH_2)_{1-10}CONR^{19}R^{20}$, $-(C_1-C_6 \text{ alkylene})-COOR^{19}$, $-CH=CH-COOR^{19}$, $-CF_3$, -CN, $-NO_2$ and halogen;

R¹⁹ and R²⁰ are independently selected from the group consisting of H, (C₁-C₆)alkyl, aryl and aryl-substituted (C₁-C₆)alkyl;

R²¹ is (C₁-C₆)alkyl, aryl or R²⁴-substituted aryl;

 R^{22} is H, (C1-C6)alkyl, aryl (C1-C6)alkyl, -C(O) R^{19} or -COOR 19 ;

 R^{23} and R^{24} are independently 1-3 groups independently selected from the group consisting of H, (C1-C6)alkyl, (C1-C6)alkoxy, -COOH, NO2, -NR¹⁹R²⁰, -OH and halogeno; and

Appellant's Brief

Application No. 10/057,323

Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

 R^{25} is H, -OH or (C₁-C₆)alkoxy.

- 84. A pharmaceutical composition for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal, comprising a therapeutically effective amount of the composition of claim 83 and a pharmaceutically acceptable carrier.
 - 86. A therapeutic combination comprising:
 - (a) a first amount of at least one peroxisome proliferator-activated receptor activator; and
- (b) a second amount of at least one sterol absorption inhibitor represented by Formula (IX):

$$Ar^1$$
— CH — Q — R_{26}
 Ar^2
(IX)

or isomers thereof, or pharmaceutically acceptable salts or solvates of the compounds of Formula (IX) or of the isomers thereof, or prodrugs of the compounds of Formula (IX) or of the isomers, salts or solvates thereof, wherein, in Formula (IX) above,

R²⁶ is selected from the group consisting of:

- a) OH;
- b) OCH₃;
- c) fluorine and
- d) chlorine;

R¹ is selected from the group consisting of

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

$$OR^5$$
 OR^4 OR^5 OR^4 OR^7 OR^7 OR^7 OR^7 OR^8 OR^8

R, R^a and R^b are independently selected from the group consisting of H, -OH, halogeno, -NH₂, azido, (C₁-C₆)alkoxy(C₁-C₆)-alkoxy and -W-R³⁰;

W is independently selected from the group consisting of -NH-C(O)-, -O-C(O)-, -O-C(O)-N(\mathbb{R}^{31})-, -NH-C(O)-N(\mathbb{R}^{31})- and -O-C(S)-N(\mathbb{R}^{31})-;

R² and R⁶ are independently selected from the group consisting of H, (C₁-C₆)alkyl, aryl and aryl(C₁-C₆)alkyl;

 R^3 , R^4 , R^5 , R^7 , R^{3a} and R^{4a} are independently selected from the group consisting of H, (C1-C6)alkyl, aryl(C1-C6)alkyl, -C(O)(C1-C6)alkyl and -C(O)aryl;

 R^{30} is independently selected form the group consisting of R^{32} -substituted T, R^{32} -substituted-T-(C₁-C₆)alkyl, R^{32} -substituted-(C₂-C₄)alkenyl, R^{32} -substituted-(C₃-C₇)cycloalkyl and R^{32} -substituted-(C₃-C₇)cycloalkyl(C₁-C₆)alkyl;

 R^{31} is independently selected from the group consisting of H and (C1-C4)alkyl;

T is independently selected from the group consisting of phenyl, furyl, thienyl, pyrrolyl, oxazolyl, isoxazolyl, thiazolyl, iosthiazolyl, benzothiazolyl, thiadiazolyl, pyrazolyl, imidazolyl and pyridyl;

R³² is independently selected from 1-3 substituents independently selected from the group consisting of H, halogeno, (C₁-C₄)alkyl, -OH, phenoxy, -CF₃, -NO₂, (C₁-C₄)alkoxy, methylenedioxy, oxo, (C₁-C₄)alkylsulfanyl, (C₁-C₄)alkylsulfinyl,

(C1-C4)alkylsulfonyl, -N(CH3)2, -C(O)-NH(C1-C4)alkyl, -C(O)-N((C1-C4)alkyl)2, -C(O)-(C1-C4)alkyl, -C(O)-(C1-C4)alkoxy and pyrrolidinylcarbonyl; or R³² is a covalent bond and R³¹, the nitrogen to which it is attached and R³² form a pyrrolidinyl, piperidinyl, N-methyl-piperazinyl, indolinyl or morpholinyl group, or a (C1-C4)alkoxycarbonyl-substituted pyrrolidinyl, piperidinyl, N-methylpiperazinyl, indolinyl or morpholinyl group;

Ar¹ is aryl, R¹⁰-substituted aryl; pyridyl, isoxazolyl, furanyl, pyrrolyl, thienyl, imidazolyl, pyrazolyl, thiazolyl, pyrazinyl, pyrimidinyl or pyridazinyl;

Ar² is aryl or R¹¹-substituted aryl;

Q is $-(CH_2)_q$ -, wherein q is 2-6, or, with the 3-position ring carbon of the azetidinone,

$$R^{12} - (R^{13})_a$$
 forms the spiro group $(R^{14})_b$; R^{12} is

 R^{13} and R^{14} are independently selected from the group consisting of -CH₂-, -CH(C₁-C₆ alkyl)-, -C(di-(C₁-C₆) alkyl), -CH=CH- and -C(C₁-C₆ alkyl)=CH-; or R^{12} together with an adjacent R^{13} , or R^{12} together with an adjacent R^{14} , form a -CH=CH- or a -CH=C(C₁-C₆ alkyl)- group;

a and b are independently 0, 1, 2 or 3, provided both are not zero; provided that when R^{13} is -CH=CH- or -C(C₁-C₆ alkyl)=CH-, a is 1; provided that when R^{14} is

-CH=CH- or -C(C₁-C₆ alkyl)=CH-, b is 1; provided that when a is 2 or 3, the R¹³'s can be the same or different; and provided that when b is 2 or 3, the R¹⁴'s can be the same or different;

 $\rm R^{10}$ and $\rm R^{11}$ are independently selected from the group consisting of 1-3 substituents independently selected from the group consisting of (C1-C6)alkyl, $\rm -OR^{19}$, $\rm -O(CO)R^{19}$, $\rm -O(CO)OR^{21}$, $\rm -O(CH_2)_{1-5}OR^{19}$, $\rm -O(CO)NR^{19}R^{20}$, $\rm -NR^{19}R^{20}$, $\rm -NR^{19}(CO)R^{20}$, $\rm -NR^{19}(CO)OR^{21}$, $\rm -NR^{19}(CO)NR^{20}R^{25}$, $\rm -NR^{19}SO_2R^{21}$, $\rm -COOR^{19}$, $\rm -CONR^{19}R^{20}$, $\rm -COR^{19}$, $\rm -SO_2NR^{19}R^{20}$, $\rm -S(O)_{0-2}R^{21}$, $\rm -O(CH_2)_{1-10}COOR^{19}$, $\rm -O(CH_2)_{1-10}CONR^{19}R^{20}$, $\rm -(C_1-C_6$ alkylene)-COOR^{19}, $\rm -CH=CH-COOR^{19}$, $\rm -CF_3$, $\rm -CN$, -NO₂ and halogen;

R¹⁹ and R²⁰ are independently selected from the group consisting of H, (C₁-C₆)alkyl, aryl and aryl-substituted (C₁-C₆)alkyl;

R²¹ is (C₁-C₆)alkyl, aryl or R²⁴-substituted aryl;

 R^{23} and R^{24} are independently 1-3 groups independently selected from the group consisting of H, (C1-C6)alkyl, (C1-C6)alkoxy, -COOH, NO₂, -NR¹⁹R²⁰, -OH and halogeno; and

$$R^{25}$$
 is H, -OH or (C₁-C₆)alkoxy,

wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal.

100. A composition comprising (a) at least one antioxidant or vitamin and (b) at least one substituted azetidinone compound or substituted β -lactam compound or isomers thereof, or pharmaceutically acceptable salts or solvates of the at least one substituted azetidinone compound or the at least one substituted β -lactam compound or of the isomers thereof, or prodrugs of the at least one substituted azetidinone compound or the at least one substituted β -lactam compound or of the isomers, salts or solvates thereof.

Appellant's Brief

Application No. 10/057,323 Paper Dated: April 18, 2006 Attorney Docket No. CV01489K

101. A therapeutic combination comprising (a) a first amount of at least one antioxidant or vitamin and (b) a second amount of at least one substituted azetidinone compound or substituted β -lactam compound or isomers thereof, or pharmaceutically acceptable salts or solvates of the at least one substituted azetidinone compound or the at least one substituted β -lactam compound or of the isomers thereof, or prodrugs of the at least one substituted azetidinone compound or the at least one substituted β -lactam compound or of the isomers, salts or solvates thereof, wherein the first amount and the second amount together comprise a therapeutically effective amount for the treatment of a vascular condition, diabetes, obesity or lowering a concentration of a sterol in plasma of a mammal.

EVIDENCE APPENDIX

None.

RELATED PROCEEDINGS APPENDIX

None.